FIG. 1A

GGAT	TGA	ACA /	AGGA	CGCA	TT T	CCCC	AGTA	C AT	CCAC	CAAC		TCC Ser			54
			TTT Phe												102
				10								15			20
			GAT Asp 25												150
			GGG Gly												198
			TTT Phe												246
														GCC Ala 85	294
					Phe					Pro				TCT Ser	342
				Trp					Ala				Phe	ACA Thr	390
			His					Gly				Ile		CTC Leu	438
		Пe					Ala				Val			TTA Leu	486

FIG. 1B

A GCC s Ala O											534
G GTG u Val											582
G AAA n Lys											630
G AAT p Asn											678
G CTG o Leu 215	Leu										726
T CGG u Arg 0											774
C ACC e Thr			V a 1								822
C ATT l Ile						Phe				Asn	870
A AGC u Ser		Ser			Ala				Glu		918
G ATG y Met 295	Thr			Asn				Ala			960

FIG. 1C

GAG AAG TTC AGA AGC CTT TTT CAC ATA GCT CTT GGC TGT AGG ATT GCC Glu Lys Phe Arg Ser Leu Phe His Ile Ala Leu Gly Cys Arg Ile Ala 310 325	1014
CCA CTC CAA AAA CCA GTG TGT GGA GGT CCA GGA GTG AGA CCA GGA AAG Pro Leu Gln Lys Pro Val Cys Gly Gly Pro Gly Val Arg Pro Gly Lys 330 335 340	1062
AAT GTG AAA GTG ACT ACA CAA GGA CTC CTC GAT GGT CGT GGA AAA GGA Asn Val Lys Val Thr Thr Gln Gly Leu Leu Asp Gly Arg Gly Lys Gly 345 350 355	1110
AAG TCA ATT GGC AGA GCC CCT GAA GCC AGT CTT CAG GAC AAA GAA GGA Lys Ser Ile Gly Arg Ala Pro Glu Ala Ser Leu Gln Asp Lys Glu Gly 360 365 370	1158
GCC TAGAGACAGA AATGACAGAT CTCTGCTTTG GAAATCACAC GTCTGGCTTC	1121
ACAGATGTGT GATTCACAGT GTGAATCTTG GTGTCTACGT TACCAGGCAG GAAGGCTGAG	1271
AGGAGAGAGA CTCCAGCTGG GTTGGAAAAC AGTATTTTCC AAACTACCTT CCAGTTCCTC	1331
ATTTTTGAAT ACAGGCATAG AGTTCAGACT TTTTTTAAAT AGTAAAAATA AAATTAAAGC	1391
TGAAAACTGC AACTTGTAAA TGTGGTAAAG AGTTAGTTTG AGTTGCTATC ATGTCAAACG	1451
TGAAAATGCT GTATTAGTCA CAGAGATAAT TCTAGCTTTG AGCTTAAGAA TTTTGAGCAG	1511
GTGGTATGTT TGGGAGACTG CTGAGTCAAC CCAATAGTTG TTGATTGGCA GGAGTTGGAA	1571
GTGTGTGATC TGTGGGCACA TTAGCCTATG TGCATGCAGC ATCTAAGTAA TGATGTCGTT	1631
TGAATCACAG TATACGCTCC ATCGCTGTCA TCTCAGCTGG ATCTCCATTC TCTCAGGCTT	1691
GCTGCCAAAA GCCTTTTGTG TTTTGTTTTG TATCATTATG AAGTCATGCG TTTAATCACA	1751
TTCGAGTGTT TCAGTGCTTC GCAGATGTCC TTGATGCTCA TATTGTTCCC TAATTTGCCA	1811
GTGGGAACTC CTAAATCAAA TTGGCTTCTA ATCAAAGCTT TTAAACCCTA TTGGTAAAGA	1871

FIG. 1D

ATGGAAGGTG GAGAAGCTCC CTGAAGTAAG CAAAGACTTT CCTCTTAGTC GAGCCAAGTT	1931
AAGAATGTTC TTATGTTGCC CAGTGTGTTT CTGATCTGAT	1991
TTCTAGAACC AGGCAACTTG GGAACTAGAC TCCCAAGCTG GACTATGGCT CTACTTTCAG	2051
GCCACATGGC TAAAGAAGGT TTCAGAAAGA AGTGGGGACA GAGCAGAACT TTCACCTTCA	211
TATATTTGTA TGATCCTAAT GAATGCATAA AATGTTAAGT TGATGGTGAT GAAATGTAAA	217
TACTGTTTTT AACAACTATG ATTTGGAAAA TAAATCAATG CTATAACTAT GTTGATAAAA	223
G	2232

FIG. 2A

CAGG	ACTO	acc 1	rgag,	ACAA	GC C	ACAA	GCT	A AC	AGA	AAA	G TG	GATT	GAAC	AAG	GACGCA	T 6	0
TTC	CCA	GTA	CATO	CAC									CGG Arg			11	. 0
															AT TAT sp Tyr	15	8
				15					2	0				2	25		
													ATT Ile 40	Gly		20	16
													GGT Gly			25	54
													AAG Lys			30)2
													GAT Asp			35	50
													AAT Asn			39	}8
									Phe				TAT Tyr 120	His		44	16
													ATC Ile			49	94
							Val					Ala	AGG Arg			54	42

FIG. 2B

			AGT Ser						590
			ATC Ile						638
			TAT Tyr						686
			TTG Leu						734
			ATC Ile 225						782
			GCA Ala				_		830
			ACT Thr						878
		Phe	GGC Gly					CAA Gln	926
			GTG Val						974
	Asn		TAT Tyr					AGG Arg	1022

FIG. 2C

TAT CTC TCG GTG TTC TTC CGA AAG CAC ATC ACC AAG CGC TTC TGC AAA Tyr Leu Ser Val Phe Phe Arg Lys His Ile Thr Lys Arg Phe Cys Lys 315 320 325 330	1070
CAA TGT CCA GTT TTC TAC AGG GAG ACA GTG GAT GGA GTG ACT TCA ACA Gln Cys Pro Val Phe Tyr Arg Glu Thr Val Asp Gly Val Thr Ser Thr 335 340 345	1118
AAC ACG CCT TCC ACT GGG GAG CAG GAA GTC TCG GCT GGT TTA Asn Thr Pro Ser Thr Gly Glu Gln Glu Val Ser Ala Gly Leu 350 355 360	1160
TAAAACGAGG AGCAGTTTGA TTGTTGTTTA TAAAGGGAGA TAACAATCTG TATATAACAA	1220
CAAACTTCAA GGGTTTGTTG AACAATAGAA ACCTGTAAAG CAGGTGCCCA GGAACCTCAG	1280
GGCTGTGTGT ACTAATACAG ACTATGTCAC CCAATGCATA TCCAACATGT GCTCAGGGAA	1340
TAATCCAGAA AAACTGTGGG TAGAGACTTT GACTCTCCAG AAAGCTCATC TCAGCTCCTG	1400
AAAAATGCCT CATTACCTTG TGCTAATCCT CTTTTTCTAG TCTTCATAAT TTCTTCACTC	1460
AATCTCTGAT TCTGTCAATG TCTTGAAATC AAGGGCCAGC TGGAGGTGAA GAAGAGAATG	1520
TGACAGGCAC AGATGAATGG GAGTGAGGGA TAGTGGGGTC AGGGCTGAGA GGAGAAGGAG	1580
GGAGACATGA GCATGGCTGA GCCTGGACAA AGACAAAGGT GAGCAAAGGG CTCACGCATT	1640
CAGCCAGGAG ATGATACTGG TCCTTAGCCC CATCTGCCAC GTGTATTTAA CCTTGAAGGG	1700
TTCACCAGGT CAGGGAGAGT TTGGGAACTG CAATAACCTG GGAGTTTTGG TGGAGTCCGA	1760
TGATTCTCTT TTGCATAAGT GCATGACATA TTTTTGCTTT ATTACAGTTT ATCTATGGCA	1820
CCCATGCACC TTACATTTGA AATCTATGAA ATATCATGCT CCATTGTTCA GATGCTTCTT	1880
AGGCCACATC CCCCTGTCTA AAAATTCAGA AAATTTTTGT TTATAAAAGA TGCATTATCT	1940
ATGATATGCT AATATATGTA TATGCAATAT AAAATTTAG	1979

FIG. 3(A)

FIG. 3(B)

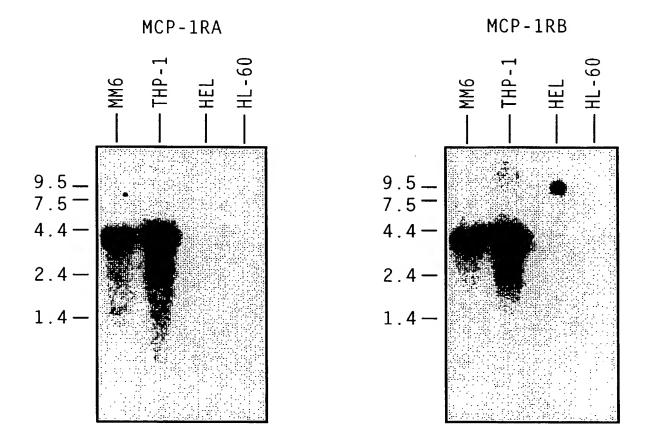
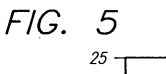


FIG.4(A)

MCP-1RA (CCR2-A) MIP-1α/RANTESR HUMSTSR IL-8RA IL-8RB	MLSTSRSRFIRNTNESGEEVTTFFDYDYGAPCHKFDVKQIGAQLLPPL METPNTTEDYDTTTEFDYGDATPCQKVNERAFGAQLLPPL MEGISIYTSDNYTEEMGS-GDYDSMK-EPCFREENANFNKIFLPTI MSNITDPQ-MWDFDDLNFTGMPPADEDYSPC-MLETETLNKYVVIIA MESDSFEDFWKGEDLSNYSYSSTLPPFLLDAAPC-EPESLEINKYFVVII	48 40 44 45 49
MCP-1RA (CCR2-A) MIP-1α/RANTESR HUMSTSR IL-8RA IL-8RB	YSLVFIFGFVGNMLVVLILINCKKLKCLTDIYLLNLAISDLLFLITLPLW YSLVFVIGLVGNILVVLVLVQYKRLKNMTSIYLLNLAISDLLFLFTLPFW YSIIFLTGIVGNGLVILVMGYQKKLRSMTDKYRLHLSVADLLFVITLPFW YALVFLLSLLGNSLVMLVILYSRVGRSVTDVYLLNLALADLLFALTLPIW YALVFLLSLLGNSLVMLVILYSRVGRSVTDVYLLNLALADLLFALTLPIW	98 90 94 95 99
MCP-1RA (CCR2-A) MIP-1α/RANTESR HUMSTSR IL-8RA IL-8RB	101 115 3 136 AH-SAANEWVFGNAMCKLFTGLYHIGYFGGIFFIILLTIDRYLAIVHAVF IDYKLKDDWVFGDAMCKILSGFYYTGLYSEIFFIILLTIDRYLAIVHAVF AV-DAVANWYFGNFLCKAVHVIYTVNLYSSVLILAFISLDRYLAIVHATN AA-SKVNGWIFGTFLCKVVSLLKEVNFYSGILLLACISVDRYLAIVHATR AA-SKVNGWIFGTFLCKVVSLLKEVNFYSGILLLACISVDRYLAIVHATR	147 140 143 144 148
MCP-1RA (CCR2-A) MIP-1α/RANTESR HUMSTSR IL-8RA IL-8RB	154 4 178 ALKARTVTFGVVTSVITMLVAVFASVPGIIFTKCQKEDSVYVCGPYFP ALRARTVTFGVITSIIIWALAILASMPGLYFSKTQWEFTHHTCSLHFPHE SQRPRKLLAEKVVYVGVWIPALLLTIPDFIFANVSEADDRYICDRFYPN- TLTQKR-HLVKFVCLGCWGLSMNLSLPFFLFRQAYHPNNSSPVCYEVLGN TLTQKRYLVKFI-CLSIWGLSLLLALPVLLFRRTVYSSNVSPACYEDMGN	195 190 192 193 197
MCP-1RA (CCR2-A) MIP-1α/RANTESR HUMSTSR IL-8RA IL-8RB	204 5 231RGWNNFHTIMRNILGLVLPLLIMVICYSGILKTLLRCRNEKKRHRAVR SLREWKLFQALKLNLFGLVLPLLVMIICYTGIIKILLRRPNEKKS-KAVRDLWVVVFQFQHIMVGLILPGIVILFCYCIIISKLSHSKGHQKR-KALK DTAKWRMVLRILPHTFGFIVPLFVMLFCYGFTLRTLFKAHMGQK-HRAMR NTANWRMLLRILPQSFGFIVPLLIMLFCYGFTLRTLFKAHMGQ-KHRAMR	243 239 239 242 246
MCP-1RA (CCR2-A) MIP-1α/RANTESR HUMSTSR IL-8RA IL-8RB	244 6 268 VIFTIMIVYFLFWTPYNIVILLNTFQEF-FGLSNCESTSQLDQATQVTET LIFVIMIIFFLFWTPYNLTILISVFQDF-LFTHECEQSRHLDLAVQVTEV TTVILILAFFACWLPYYIGISIDSFILLEIIKQGCEFENTVHKWISITEA VIFAVVLIFLLCWLPYNLVLLADTLMRTQVIQETCERRNNIGRALDATEI VIFAVVLIFLLCWLPYNLVLLADTLMRTQVIQETCERRNHIDRALDATEI	292 288 289 292 296

FIG. 4(B)

	<u>295 7 313</u>	
MCP-1RA (CCR2-A)	LGMTHCCINPIIYAFVGEKFRŠLFHIALGCRIAPLQKPVCGGPGWRPGKN	342
MIP-1α/RANTESR	IAYTHCOVNPVIYAFVGERFRKYLRQLFHRRVAVHLVKW	327
HUMSTSR	LAFFHCCLNPILYAFLGAKFKTSAQHALLTSWSRGSS	325
IL-8RA	LGFLHSCLNPIIYAFIGONFRHGFLKILAMHGLVS	327
IL-8RB	LGILHSCLUPLIYAFIGOKFRHGLLKILAIHGLIS	331
·		
MCP-1RA (CCR2-A)	VKVTTQGL[]DGRGKGKSIGRAPEAS[JQDKEGA	374
MIP-1α/RANTESR	LPFLSVDRLE-RVSSTS-PSTGEHELLSAGF	359
HUMSTSR	LKILSKGKRGGHSSVSTESESSSFHSS	35
IL-8RA	KEFLARHRVTSYT-SSSVNVSSNL	350
IL-8RB	KDSLPKDSRPSFVG-SSSGHTSTTL	35



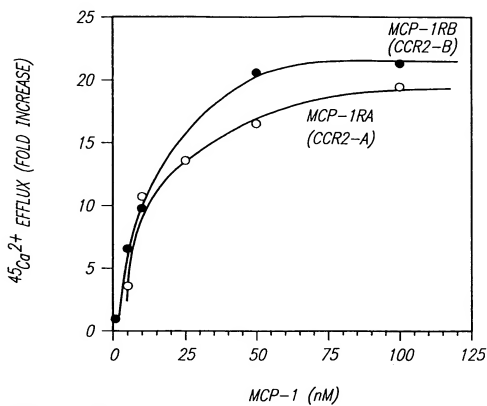
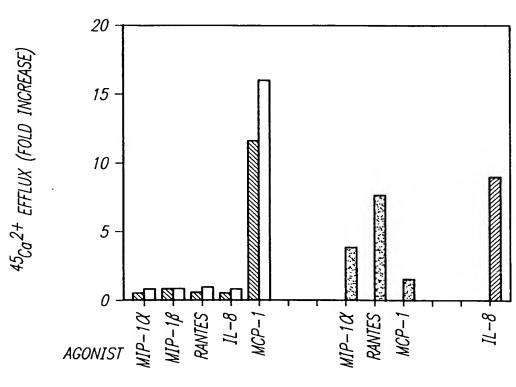


FIG. 6



MCP-1RA

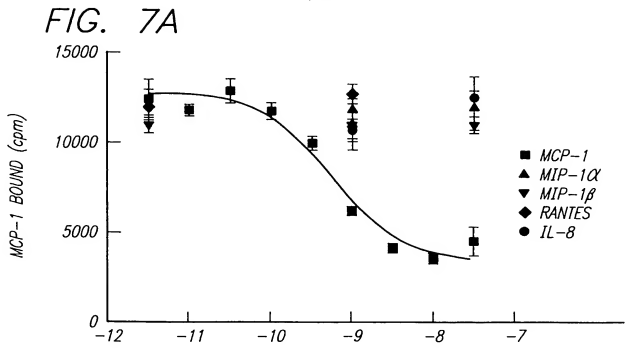
MCP-1RB

cRNA

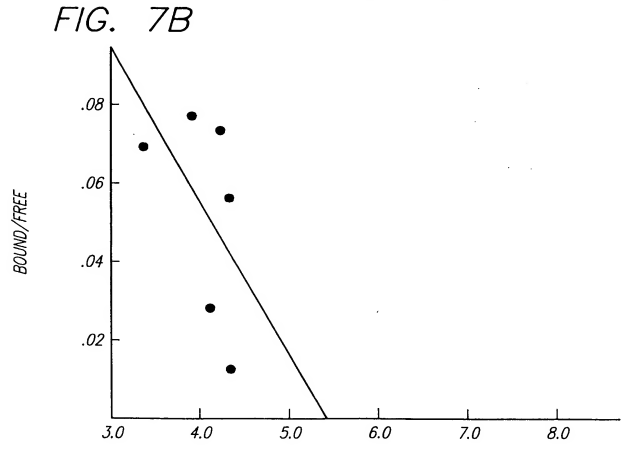
MCP-10/RANTES R

IL-8RA





LOG [CHEMOKINE] (M)



MCP-1 BOUND (M, $X10^{-11}$)

